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- ② Inventor: Sajic, Branko
- W Hair conditioner compositions containing fatty acid ester derivatives of alkanolamines.
- The present invention provides hair conditioner compositions comprising an effective conditioning amount of an alkanolamine derivative selected from the group consisting of: an alkanolamine sett of formula it:

#### where

A and B are are the same or different and represent lower alkylene having 1-8 carbon atoms; and R; and R; are the same or different and represent alkyl or alkenyl groups having 8 to 40 carbon atoms; an alkanolamine ester quaterney ammonium set of formula III:

where

- R is lower alkyl having 1-6 carbon atoms;
- A and B are the same or different and represent lower alkylene having 1-8 carbon atoms; and
- $R_1$  and  $R_2$  are the same or different and represent alkyl or alkenyl groups having 8 to 40 carbon atoms; and X is Cl<sup>-</sup>, Br<sup>-</sup>, I<sup>-</sup>, RSO<sub>3</sub><sup>-</sup>, or RSO<sub>4</sub><sup>-</sup>; and
- mixtures of the alkanolamine ester and the alkanolamine ester quaternary ammonium salt.
- The invention further provides conditioning compositions comprising alkanolemine derivatives of formulas it and/or it and silicone conditioning compounds. Methods for preparing such conditioning compositions are also disclosed.

#### BACKGROUND OF THE INVENTION

#### Field of the Invention

This invention relates to hair conditioning compositions and more specifically it relates to hair conditioning compositions comprising an askanolamine derivative conditioning compound. It also relates to conditioner compositions comprising a silicone oil conditioning compound and an alkanolamina derivative.

#### Description of the Related Art

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Quaternary ammonium compounds have been extensively used as conditioning agents in both rinse-out and leave-on heir conditioners. In addition, long chain alkyl amines and alkanolamines have routinely been incorporated into hair conditioner compositions for use as emulsifiers, suspending or thickening agents. The quarternary compounds mentioned above are well known to provide a conditioning effect on heir by improving detangling, wet combing, dry combing, wet and dry feel and static/fly away. However, quaternary ammonium salts alone are unable to provide sufficient conditioning effects to hair conditioning and especially to hair that is chemically treated, long, or naturally wavy.

For that reason, silicone oil conditioning compounds have been utilized to enhance the performance of hair conditioning compositions. However, silicones are not compatible with all quaternary salts or all zo conditioner bases. For example, most low molecular weight (low vapor pressure) and high molecular weight (high vapor pressure) silicone compounds, when combined in a conditioner with a di- or tri-long-chain alkyl quaternary ammonium salt, do not yield good conditioning effects on hair. Such conditioners tend to result in an oily feel and poor overall combing properties. See, for example, U.S. Patent No. 4,777,037.

Quaternary ammonium salts have routinely been incorporated into fabric softening compositions. For 25 example, alkanolamine ester quaternary ammonium saits of the general formula (

where

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R is methyl;

A and B are athylene;

and R<sub>1</sub> and R<sub>2</sub> are alkyl or alkenyl groups having 14 to 18 carbon atoms;

have been incorporated into various laundry fabric softening compositions. In particular, U.S. Patent No. 3,915,867 disploses the use of N-methyl, N,N-di-(\$-C12-acyloxyethyl),N-\$-hydroxy ethyl ammonlum 45 methylsulfate as the active softening ingredient in a fabric softening composition.

The currently utilized quaternary ammonium salts are frequently made from synthetic sources, are normally not biodegradable, and have a high toxicity towards fish and water plants.

Similar quaternary ammonium compounds have also been incorporated into fabric soiteners. See, for example, European Patent application No. EP 295 386 A2; European Patent Application No. EP 370 675 A2; so J.A.C.S., 104: 458-61 (1982); and Comun. Jorn. Com. Esp. Deterg. 20: 181-192 (1989).

Silicone oil compounds traditionally have been incorporated into hair conditioning compositions with various quaternary ammonium saits. Exemplary of the quaternary ammonium saits that have been incorporated into silicone containing conditions are: diffrydrogenated) tallow dimethyl ammonium chloride; coco bis(hydroxyethyl) methyl ammonium chloride; trimethyl soyaalkyl ammonium chloride; cocoalkyl ss trimethyl ammonium chlorida; di-C12-C18-alkyl dimethyl ammonium saits; dicocoalkyl dimethyl ammonium chloride; trimethyl tallow ammonium chloride; lauryl trimethyl ammonium chloride; tri-Ce-se alkyl methyl ammonium chlorides; and di-C14-15-alkyl dimethyl ammonium chloride salts.

Representative hair conditioner compositions are disclosed in U.S. Patent No. 4,777,037. That patent discloses hair conditioning compositions which comprise a di-Ct<sub>2</sub>-1s-sikyl-di-Ct<sub>2</sub>-2-sikyl ammonium saft and a polydimethyl sitoxane (a volatile cyclic sificone).

Accordingly, conditioning compositions are desired that employ a conditioning compound or allicons is and a conditioning compound where the conditioning compound is compatible with the allicone as the prepared from a natural resource, readily and ultimately biodegradable, and only minimatly toxic to fish and water plants.

#### SUMMARY OF THE INVENTION

The present invention provides hair conditioner compositions comprising fatty acid ester derivatives of alkanolamines which impart excellent conditioning properties to hair.

The present invention provides hair conditioners comprising an effective conditioning amount of an alkanolamine derivative selected from the group consisting of:

16 an alkangiamine ester of formula it:

#### where

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A and B are are the same or different and represent lower alkylene having 1-8 carbon atoms; and R; and Re are the same or different and represent airly or alkonyl groups having 8 to 40 carbon atoms; an alkenolamine ester quaternary ammonfulum sat of formula filt:

#### where

R is lower alkyl having 1-6 carbon atoms;

A and B are the same or different and represent lower alkylene having 1-6 carbon atoms; and 8, and 8e are the same or different and represent alkyl or alkenyl groups having 8 to 40 carbon atoms; and

X is Cit., Bit., It., RSO<sub>3</sub>T, or RSO<sub>4</sub>T; and

mbdures of the alkanolamine ester and the alkanolamine ester quaternary ammonium salt. The present invention turther provides hair conditioning compositions comprising a silicone oil concitioning compound and an alkanolamine derivative of Formula II and/or Formula III

The present invention also provides methods for preparing hair conditioning compositions comprising a compound of formula it and/or till and a stiticone oil conditioning agent.

It has been surprisingly discovered that the alkanolamino derivatives of formulas il and fill are capable of functioning efficiently in combination with sifecone conditioning agents of various vapor pressures and molecular weights.

# DETAILED DESCRIPTION OF THE INVENTION

It has been unexpectedly discovered that when hair conditioning compositions are prepared to contein a tatty acid exter derivative of an altenotamine or a silicence oil conditioning composition and a fatty acid exter certification of an altenotamine, the resultant heir conditioning compositions provide excellent conditioning effects on heir.

Thus, the present invention provides hair conditioners comprising an effective conditioning amount of an alkanolamine derivative selected from the group consisting of:

an alkanolamine ester of formula II:

where
A and B are are the same or different and represent lower alkylene having 1-6 carbon atoms; and

R<sub>1</sub> and R<sub>2</sub> are the same or different and represent alkyl or alkenyl groups having 8 to 40 carbon atoms; an alkanolamine ester quaternary ammonium salt of formula III:

where

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R is lower alkyl having 1-6 carbon atoms;

A and B are the same or different and represent lower alkyleng traving 1-6 carbon atoms; and

R<sub>1</sub> and R<sub>2</sub> are the same or different and represent alkyl or alkenyl groups having B to 40 carbon atoms; and

X is CIT, BrT, tT, RSO<sub>3</sub>T, or RSO<sub>4</sub>T; and

mixtures of the alkanolamine ester and the alkanolamine ester quaternary ammonium salt.

In addition, the present invention encompasses hair conditioning compositions comprising effective conditioning amounts of a compound of formula II and/or formula III and a silicone oil conditioning

in the hair conditioning compositions of the passent invention, these alkanolamine derivatives function above or in undorn with sitione oil conditioning companets to provide conditioning effects on his responsibility of the standard of the standard control of the standa

The hair conditioning compositions prepared to contain both an alkanolarimat derivative of formula II so and/or III and a silicone contilioring compound were also unexpectedly lound to exhibit conditioning properties greater than the aggregate of the conditioning properties of each individual component. In addition, the compositions of the present invention are prepared using alkanolarimic derivatives that demonstrate minimal fish and water plant toxicity, are readily and utilizately biodegradable and are

prepared from simple natural resources.

The present invention relates to hair conditioning compositions which contain water and an alisonalaming desirative conditioning agent. These alternatives have the chemical formulas if and ill shown above. The invention also relates to hair conditioning compositions which contain water and a combination of a silicone and an alternativemic devirative of formula tile rill. The compositions of this invention are useful both as water dispersions or aquecus emulsions preferably stable emulsions in which water compositions the external charges.

Those skilled in the art will recognize a variety of synthetic memodologies for preparing the alkanolenine devicefree of formulas it and III. Exemplary of suitable preparative methods for synthesizing an 10 alkanolenine ester of formula! It is an exid catalyzed destriction of a fatty solid with an alkanol. In addition, such an alkanolenine ester may be prepared by a base-catalyzed transesterflication of a methyl ester in the presence of an alkanolenine.

The resulting atkanolamine esters may be quaternized using any suitable quaternizing agent, such as for example, dimethyl sulfate.

A synthesis of compounds of these classes is found in U.S. Patent No. 3,915,867.

The silicone compounds which may be used in the hair conditioning compositions of the present invention are well known and include those that have been generally sught to be useful in a variety of emulsions. The silicones useful for incorporation linto the conditioning compositions of the invention are silicone fulds as well as silicone gums. Such silicones will have visuosities ranging from a few fundred so centropies to about 1,000,000 centropies. A bland of high and low silicones can be utilized in order to obtain the desired conditioning effect and to facilities incorporation line in antic conditione procust.

Among the silicone compounds sutitable for incorporation into conditioning compositions of the invention are dimethicones and cyclomethicones which may be represented by the formula:

wherein R is a 1 to 3 curbon alkyl group, n is a linteger from 3 to 10, preferably from 3 to 7, and the unsatilified veheces on the oxygion and silicon aims at the ends of the chain may be joined to one enother to form a cyclic structure. Sulfablis silicone compounds are, for exemple, U.C.C. Y-7207, sold by Union 3 cachide Corporation in which each R is methyl and which typically comprises by weight 99.4% betarner, 0.0% infiner and traces of the pontamor and hexamir, and SWS-0.0314. sold by SWS Silicones. a Division of Sautifer Chemical Company, in which R is mothly and which is sustainatily, all terramer; and Dev Corning, 444 fluid, sold by Dow Corning, i.e., in which R is methyl and which typically comprises by weight about 89% internance, about 11.8% penhamer and traces of trimer and hexamer. Typical vegor pressures of 40 silicones are shown in the table below. These vapor pressures were determined using Dow Corning 344 fluid at various temperatures.

Temperature	Vapor Pressure, mm Hg
20C	1
64 ° C	10
77 · C	20
92°C	40
101 °C	60
114°C	100
155 °C	400
178°C	760

Such silicones are said by one manufacturer to be useful in various coernetic compositions such as antiperspirants, deoderants, has prayes, hair cottoring and hair growning products, and because of their low viscosity and surface tension provide a light sifey feel on hair and skin. These silicones, are also reported to be non-prayes but to provide suitble labrication.

Also suitable for use in the invention are amodimethicones of the formula:

$$HO = \begin{bmatrix} GH_3 \\ Si-O \\ GH_3 \end{bmatrix}_X \begin{bmatrix} OH \\ Si-O \\ GH_2 \end{bmatrix}_y$$

$$GH_2 \\ GH_2$$

$$GH_3 \\ GH_4$$

$$GH_4 \\ GH_5$$

$$GH_5$$

$$GH_5$$

$$GH_6$$

$$G$$

where x is an integer at least equal to 4, and y is an integer at least equal to 4.

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Such silicone compounds may be synthesized to have molecular weights and vapor pressures that range from low to high. In addition, such amodimethicanes may be quaterrized to yield cationic amine functional 15 polymer emulsions. Representative emulsions are available from Dow Corning and include 929 cationic emulsion, Silicone 02×7224, and Silicone DC X2-9839.

The preferred compositions of the invention typically include water, an alkanolamine derivative of formula II and/or III. Water typically constitutes at least about 70 weight percent of the weight of the conditioning composition, and more preferably about 80 weight percent.

The silicones may be present in the hair conditioning compositions of this invention in amounts of from about 0.5 to about 15 weight percent of the composition, Preferably, these silicones are present at from about 1 to about 10 percent by weight of the conditioning composition. Particularly preferred amounts of silicone in the compositions of the invention are from about 1.0 to about 8 percent of the composition.

The alkanolamine derivative conditioning agents are preferably present at from about 0.5 to about 20 so percent by weight of the composition as an active ingredient. More preferably, the alkanolamine derivative conditioning agent is present at from about 1 to about 5 weight percent, as an active ingredient, in particularly preferred compositions, alkanolamine derivative is present at from about 1-3 percent of the

composition.

In certain compositions according to the invention, the alkanolamine derivative is present as a mixture of 30 an alkanolamine ester and alkanolamine ester quaterary ammonium salt. In such compositions, the combined amount of the alkanolamine ester alkanolamine ester alkanolamine ester alkanolamine ester dusterary ammonium salt ranges from about 0.1% to about 2.0% of the conditioner composition. In these compositions, the ratio of alkanolamine ester to alkanolamine ester quaterary ammonium salt ranges from about 1.19% to about 9.11. In more perfered compositions, the ratio of alkanolamine ester to alkanolamine ester quaterary ammonium salt ranges and the compositions.

In certain embodiments of the invention, the alkanolamine derivative is introduced into the composition as an about 50-88% milant of the alkanolamine derivative in abonit or glycol. In certain prefer embodiments, the alkanolamine derivative conditioning compound is introduced as an about 70-80% milature of the alkanolamine derivative in the alkanolamine derivative interfered embodiments of the 40 invention are prepared using a milature of the alkanolamine derivative in propytene glycol, ethanol or isoscropanoli.

Proferred elkanolamino derivatives according to the invention are N-methyl, NN-diff-partially hydrogenated talloyloxyethyl, Ny-d-hydroxyethyl ammonium methyl suffate; and N-Methyl, NN-diff-partially hydrogenated partingloxyethyl), Ny-dhydroxyethyl ammonium methyl suffate.

To noted that the long aliphatic chains of the the quaternary ammonium conditioning agents, designated by R, and R) in formulas if and Ills above, need not be solely or printingly of one chain length; i.e., the long chain need not be celly, myristyl, lauryl or stearyl. Rather, conditioning agents having long aliphatic chains R, and Rp containing a mixture of lengths can be used. Such conditioning agents are conveniently prepared from naturally cocurring materials, such as tallow, coconut cit, soys cit and the like.

or from yortholically produced mixtures.

The compositions of this invention containing only water, sitions and the alkanolarnine derivative are milky-white, relatively viscous disposition. Shows be compositions are stable to playes separation at a temperatures of about 0 for about 45 °C for at least 30 days after their preparation, and are typically stable to shake separation indefinitely a five Profession 50 °C from about 20 to 25 °C.

The compositions of this invention can also be in the form of emulsions that contain additional amounts of hydrophics andorf hydrophics ceruitalities. Fundisons containing additional emulsition materials are particularly preferred. It is preferred that those emulsions be stable to phase separation at a temperature of about 45°C for a period of about 30 days after their preparation. The emulsions are more preferably stable

to phase separation at temperature normally found in commercial product storage and shipping for periods of six months or more.

The compositions of the invention may also contain trickeners or emulsifiers. The thickeners and emulsifiers may be long chain tatty abcorbs having from about 11 to about 22 carbon atoms. These aborbits can be used alone, or in admixing with each other. When included in the compositions, the abcorbit is preferably present at from about 0.5 to about 10 weight percent of the composition, and more embersable at from about 2 to about 5 weight percent.

Lauryl alcohol, Jeryl sacholi, myristyl alcohol, cethy salcohol, steamyl alcohol, and the like, and mixtures thereof are contemplated herein. In addition, mixtures of natural or synthesis fasty alcohols having fasty chains of from about 11 to about 22 carbons are also useful. Several such mixtures are available commercially, and are exemplified by the material containing a mixture of synthesis about with 12 to 15 carbons in the algri chain sold under the trademark NCDODL 25 Needl Chemical Company, and the material containing a mixture of synthesis alcohols wish chain lengths of 12 to 16 carbons sold under the trademark ALPOL 1216 Alcohol by Concor Chemically.

Fatty alcohols of the above discussed carbon chain lengths which are ethoxylated to contain an average of one or two moles of ethylace oxide per mole of talty alcohol can be used in place of the fathy alcohols immensives. Exemplace of auch useful ethoxylated fathy acods include ethylane glycol oxyl other, polyoxylately, or place of the place of the fathy and the fathy and the fathy alcohols contained (2) steary) ether, and the file, the exemplary compounds having CTFA Dictionary names of Carletin and Standberg. Careactivities.

The thicknews and emulsifiers suitable for use in the invention may also be compounds such as, for sxample, hydroxysthyloellulose (available under the tradename Natroscile from Aqualion), hydroxysthyloellulose (available under the tradename Natrocole from Dow Chemical), xantha gum (available under the tradename Natrocole from Dow Chemical), xantha gum (available under the tradename Natrocole and National National (available under the tradename Natrocole and National National (available under the tradename Natrocole and National National

under the tradename Lexamine® S-13 from Inclass).

A tentiary unidcamine can also be present in the hair conditioning compositions of this invention, and is present in particularly preferred compositions at a concentration of from about 0.1 to about 2 weight percent of the composition, and more preferably at from about 0.25 to about 1 weight percent.

The triting ambicamines useful herein have structures conforming to the formula R<sup>-</sup>(C/e O)N-HR<sup>-</sup>A-Figh-werein R<sup>-</sup> is a fatty other shoring about 11 to about 17 cations strone, R<sup>-</sup> is an eligible group lavaring 2 or 3 carbon stores and each RP is eithyl or methyl. Exemplary, useful, totary ambidamines include dimethylaminecopy at seammide, delictlylaminecityl setemplica and enarthylaminecopy at seammide, delictlylaminecitylaminecopy at seammide, and enarthylaminecopy myristamide. The R<sup>-</sup> group of the tertiary ambidamines can also be prepared from coccenst, soys and tailow fatty socids, or the like.

The hair conditioning compositions of this invention suitably are near neutral to sligntly acidic in pH 39 value. Thus, the hair conditioners of this invention preferably have pH values of from about 3 to about 8, and more preferably thom about 3.5 to about 6.0

Ingredients in addition to water and the previously discussed ingredients can also be present in the composition of the invention. These additional impredients include, but are not infriend to, polyhydrid and provides and the invention. These additional impredients include, but are not infriend to, polyhydrid active and the interest of the invention of the interest of the interes

The compositions of the invention provided excellent results in the erreas of desangling, well- and discombability, wet and dry feet, statistifysaws, sheen, body etc. Three conditioning proporties were
demonstrated with conditioner compositions incorporating a variety of sticcine compounds having a wide
for engling of molecular weights and vapur pressures. These results were demonstrated in each tests and in
shortstry tests on evacities of hair. As pointed ut above, the later conditioning compositions prepared to
contain both an elizandamine derivative of formula it and/or it and a sticcine conditioning composition or
unexpectedly found to exhibit conditioning propriets greater than the aggregate of the conditioning

properties of each individual component.

The conditioning compositions of the present invention are readily manufactured using any conventional emulsification process. The manufacture of the conditioners may be effected using either single-phase hot processes or multi-phase processes.

One skilled in the art will recognize that modifications may be made in the present invention without overlating from the spirit or scope of the invention. The invention is illustrated thather by the following examples which are not to be construed as limiting the invention or scope of the specific procedures described herein.

# 10 Example 1

Preparation of N-methyl, N, N-di (8-partially hydrogenated talloyloxyethyl), N-8-hydroxy ethyl ammonium methyl sulfate

# 15 1. N,N-di(β-partially hydrogenated talioyloxyethyl), N-β-hydroxy ethylamine

To a suitable reaction flask equipped with a stirrer, nitrogen inlet and thermometer was added 61.4 g of partially hydrogenated sallow faitly add. The atmosphere in the flask was purged with nitrogen, after which prosphoric add (thyPD., 0.12g) and triethanolamine (TEA, 24.4g) were added. The reaction flask was heated at about 155-160°C while removing water until the reaction was complete.

The resulting N,N-di(s-partially hydrogenated talloyloxyethyl), N-s-hydroxy ethylamine may be incorporated into conditioner compositions or may be quaternized as described below. If it is used as the free amine, it can be added nest or as a mixture of the amine in an alcohol or glycol.

#### 25 2. Quaternization with dimethyl sulfate

To 100g of the diester amine product prepared have in part 1 was added 18.1g of dimethyl suifate and the minuture heated for about 2 hours at 87°C. A Viscous dester quaternary ammonium dimethyls sulfate product was obtained. Prior to incorporation into a conditioner composition, the diester quaternary ammonium methyl sulfate product was mixed with isopropanol, ethanol or propylere glycot to yield an about 50°. 98% mixture of the quaternary ammonium methyl sulfate in isopropanol, ethanol, or propylere glycot. The diester quaternary ammonium methyl sulfate in isopropanol, ethanol, or propylere glycot. The diester quaternary ammonium methyl sulfate in sopropanol, ethanol, or propylere glycot. The diester quaternary ammonium methyl sulfate product prepared according to this procedure may also contain triebrandamiem and ordinester amine.

## 35 Example 2

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Preparation of N-Methyl, N.N-di(8-partially hydrogenated palmitoyloxy ethyl), N-g-hydroxyethyl ammonium methyl sulfate

# 40 1. N,N-di(β-partially hydrogenated palmitoyloxy ethyl), N-β-hydroxy ethylamine

A 4-neck, 1 liter reaction flash fitted with a stirrer, theremoneter, nitrogen lotet, and a vecuum distillation assembly was charged with 1.78 moiss of partially hydrogenated palm fetty acid methyl sets; 133.2q (0.893 moiss) of TEA and 8p of a 25% solution of sodium methods. The mixture was stirred and heated to 45 Pit-105\*C under vacuum (about 28 inches of Hg) and nitrogen flow (about 5 \* ccmin) for about 1.75 hours. 579a of resulting anime ester was obtained.

The resulting NA-dif-pertially hydrogenated psimitoyloxy ethyl). N-p-hydroxy ethylamine may be incorporated into conditioner compositions or may be quaernized as described below. If it is used as the fine armie, it can be added earl or as a mixture of the armine in an alcohol or glycol.

# 2. Quaternization with dimethyl sulfate

The diester amine obtained in part 1 of Example 2 above was quaternized with dimethyl sulfate using substantially the same procedure as set forth in part 2 of Example 1. Prior to use in a conditioner composition, he diester quaternary ammonium methyl sulfate product was mixed with isopropano), othanol or propylene glycol to yield an about 59-65% mixture of diester quaternary ammonium methyl sulfate in isopropanol, ethanol, or propylene glycol. The diester quaternary ammonium methyl sulfate product prepared according to this procedure may set to contain trieflanodemine and/or diester amine.

#### Example 3

#### Preparation of a Rinse-Out Hair Conditioner Composition

To a suitable vessel equipped with mixing, heating, and cooling capabilities were added about 468g of deionized water. The water was then mixed as 3.0g of hydroxyethylcollulose was added to the vessel. This mixture was mixed for about 5 minutes after which about 0.05g of a 50% aqueous solution of NaOH was added and the resulting mixture mixed until smooth and free of lumps. To this mixture was then added about 6.0g of N-methyl, N,N-citig-partially hydrogenated talloyloxyethyl), N-g-hydroxy ethyl ammonium nethyl sulfate as a 75% mixture in propylene glycol, and the resultant mixture heated to 79-75°C. To the 70-75 °C mixture was added about 10g cetearyl alcohol. After mixing for about 30 minutes, the mixture was cooled to about 45°C, and 12.5g of a 20 percent aqueous patassium chloride solution added. The pH was adjusted to between 3.5 and 4.0 with 50 percent aqueous citric acid or 50 percent aqueous NaOH. The resulting conditioner composition, 3A, had a pH of 4.0.

Conditioner compositions 3B and 3C were prepared using essentially the same procedures set furth above for preparing composition 3A. Table I below shows the amounts of each component in compositions 3A, 3B and 3C. Each of these compositions resulted in excellent conditioning of heir as demonstrated in both salon and laboratory settings.

Conditioner compositions 3A, 3B and 3C demonstrated excellent stability at 20 °C for three months, and on at 2 °C and 45 °C for 30 days.

TABLE 1

Component'	3A	3B	3C
delonized water	93.56	93.46	74.89
hydroxyethyl cellulosa	0.60	0.70	0.60
sodium hydroxide (NaOH)	0.01	0.01	0.01
tallow ester gusternary ammonium sait?	1.33	-	-
palm aster quaternary ammonium salt <sup>3</sup>	-	1.33	20.0
cetearyl alcohol	2.00	2.00	2.0
potassium chloride (20% aqueous solution)	2.50	2.50	2.50
citric acid (50% aqueous solution)	Q.S.	Q.S.	Q.S.
sodium hydorxide (50% aqueous solution)	Q.S.	Q.S.	Q.S.
Ha	4.0	4.0	4.0

1 component amounts are expressed as percent of final composition 2 N-methyl, N,N-di(β-partially hydrogenated talloyloxyethyl), N-β-hydroxyethyl ammonium methyl sultate (75% in propylene glycol)

3 N-methyl, N,N-di(g-partially hydrogenated palmitoyioxy ethyl), N-g-hydroxy

ethyl ammonium methyl suifate (75% in propylene glycol)

# Example 4

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## Preparation of a Hair Conditioner Composition

To a suitable vessel equipped with stirring, heating, and cooling capabilities was added 454g of deionized water, 2.5g of N,N-dimethyl N-stearyl amine, and 6.5g of a 50% aqueous solution of citric sold. 50 This mixture was stirred and heated to 70-75 °C. At 70-75 °C, 15g of cetyl alcohol and 11g of a 70 percent mixture of N-methyl, N, N-di(8-partially hydrogenated talloyloxyethyl), N-8-hydroxy ethyl ammonium methyl sultate in propylene glycol were added and the mixture emulsified for 20-30 minutes at 70-75 °C. The mixture was then cooled to about 50 °C at which time 6.5g of a 20 percent agueous potassium chlorido (KCI) solution was added. This mixture was then stirred well and 10g of Silicone DC 345 added. The pH ss was checked and adjusted if necessary to between 3.5 and 6 with 50% aqueous clinic acid or 50 percent aqueous NaOH. The resulting conditioner composition, 4A, had a final pH of 3.6.

Conditioner compositions 4B, 4C, 4D, 4E, 4F, and 4G were prepared using essentially the same procedure set forth above for preparing composition 4A. Table II below shows the amount of each

component in compositions 4A-4G. Each of these compositions resulted in excellent conditioning of hair as demonstrated in both salon and laboratory settings.

Table I

40 4C 4D 48 46 Component<sup>3</sup> ΔA 63 43 89 44 89.84 91.34 90.44 92.54 Deionized water 90 44 0.50 0.50 0.50 0.50 0.50 0.50 N,N-dimethyl, N-stearyl amine 0.50 0.20 0.10 0.10 0.10 0.10 0.10 50% aq. citric acid tallow ester quaternary ammonium salt2 2.66 2.66 2.66 2 66 2.66 2.66 2 88 palm ester quaternary ammonium salt<sup>3</sup> 3.00 3.00 3.00 3.00 2.50 3.00 3.00 cetyl alcohol 1.30 1.30 2.40 1.30 20% someous KCI 1.30 1.30 2.40 2.00 2.00 2.00 2.00 Silicone DC 345 2.00 Q.S. Q.S. a s Q.S. Q.S. O.S. 50% ag, NaOH 08 OS o.s. Q.S. Q.S. Q.S. Q.S. 50% ag, citric acid as. 0.5 Brij® 7214 Bril 585 0.5 1.00 Promulgen® G<sup>6</sup> 5.71 Silicone Q2 72247 (35%) 0.20 DMDM hydantoin 50 3.8 5.0 4.0 4.5 3.6 3 89

- I component amounts are expressed as percent of final composition
- 2 N-methyl, N.N-di(β-partially hydrogenated talloyloxyethyl), N-β-hydroxyethyl ammonium methyl sulfate (75% in propylene glycol)
  - 3 N-methyl, N.N-di(\$-partially hydrogenated palmitcyloxy ethyl), N-\$-hydroxy ethyl ammonium methyl suffate (75% in propylene glycol)
  - 4 CH<sub>3</sub>(CH<sub>2</sub>)<sub>14</sub>CH<sub>2</sub>(OCH<sub>2</sub>CH<sub>2</sub>)<sub>2</sub>: OH, Brij is a registered trademark of ICI Americas
  - 5 CH<sub>2</sub>(CH<sub>2</sub>)<sub>14</sub> CH<sub>2</sub>(OCH<sub>2</sub>CH<sub>2</sub>)<sub>20</sub>OH, Brlj is a registered trademark of ICI Americas
  - 6 mixture of cetyl/stearyl alcohols and polyoxyethylene ethers of cetyl/stearyl alcohol. Promulgen is a registered trademark of Amerchol Corp.
  - 7 cationic emulsion of noncrosslinking, amine functional polymer (Dow Corning)

### Example 5

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# Preparation of a Hair Conditionar Composition

To a suitable vessel equipped with stirring, heating, and cooling capabilities was added 196g of delorized water and 1.0g of N.N-dimethyl.N-stearyl emine. The niciture was heaten with stirring to about 70 °C. Alter all the anime healted, 1.2g of N-Mathyl, N, N-d (6-partially hydrogenested palmitoploxy entry), N-d-hydrogenesty ammonium methyl sultate was added with stirring and the mixture subsequently codel of the about 30 °C. At 30 °C. 2g of Sillicone DC 345 was added and the mixture stirred sufficiently to produce an emulsion. The pH was checked and adjusted to between 4 and 5 with 50 percent aq. ciric acid or 50 sq. percent NaOH. The resulting conditioner product, 5A, net a pH of 48.

Conditioner compositions 5B and SC were prepared essentially according to the procedure set torth above for preparing composition 5A. Table III below shows the amount of each component in compositions 55 5A-SC. Each of these compositions resulted in excellent conditioning of hair as demonstrated in both salon and laboratory settings.

Table III

Component	5A	58	5C
delonized water	97.90	97.95	96.16
N,N-dimethyl,N-stearyl amine	9.50	0.50	9.50
palm ester quaternary ammonium sait?	0.60	0.55	0.55
silicons DC 345	1.00	1.00	-
50% eq. citric acid	Q.S.	Q.S.	Q.S.
50% aq. NaOH	Q.S.	Q.S.	0.8.
silicone Q2-7224 (35%)	-		2.85
pH	3.5	3.5	3.5

1 component amounts are expressed as percent of final composition 2 N-methyl, N, N-di (£-partially hydrogenated palmitoyloxy ethyl), N-£-hydroxy ethyl ammonium methyl sulfate (75% in propylene glycol)

## Example 6

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# Preparation of a Hair Conditioner Composition

To a suitable vessel equipped with healting, string, and cooling capabilities and chapped with 439 of described were was added 2.59 of N-M-Greenty N-Massaylamina, and 11.0 g of N-melty, N-M-Sigh-partially 21 hydrogenated talloylovyethyl. N-Sigh-partially atmosphare methyl suitable while healting to about 70°. 25°. O. Al between 70° and 75°. O. 12 50° of total bottoch was added and the mitruse testine for 50° minuse. The mitrure was then cooled and et about 50° C 65° g if e 20° percont aqueous solution of KD was added, followed by 10° og of Silicence Of 30° de 30°

Compositions 8A - 8I were prepared essentially according to the method set forth above to prepare composition 8A; these compositions are shown below in Table IV. Each of these compositions resulted in excellent conditioning of hair as demonstrated in both state and laboratory settings.

45	40	35	30		25	20	15	10	5	
Component,		₽ÿ	89	Jab 26	Table IV 60	39	Æ	99	핆	a
deionized water		85.33	84.83	84.63	90.54	84.83	93.04	86.83	83.38	86.83
N,N-dimethyl, N-stearylamine	damine	0.50	0.50	0.50	0.50	0.50	t	0.50	0.75	0.50
taliow ester quaternary ammonium salti		2.66	1	;	:	ŧ	١.	1	1	;
paim ester quaternary ammonium salt <sup>3</sup>		;	2.66	2.66	2.66	2.66	2.66	2.66	2.66	2.66
ceryi alcohol		2.50	3.00	3.00	3.00	3.00	3.00	3.00	4.00	3.00
20% aqueous KCl		1.30	1.30	1,30	1.30	1.30	1.30	1.30	1.30	1.30
Silicone DC 345		2.00	2.00	2.00	2.00	2.00	;	;	2.00	;
Silicons Q2-7224		5.71	5.71	5.71	1	ı	;	:	;	5.71
Cationic Emulsion 929*	*	ı	,	;	:	5.71	1	;	5.71	;
DC X2-8939*		1	:	ŧ	;	;	t	5.71	:	:
citric acid		O.S.	0.5	0.5	a.s.	0.5	0.5	0.5	0.5	o,s
aq. NaOH		0.5.	0.8.	0.5.	0.5	0.5	0.5.	Q.S.	Q.S.	0.5
DMDM Hydantoin		:	;	0.2	;	1	;	:	0.2	i
Ha.		4.0	4.0	4.0	4.0	4.0	4.0	4.0	4.0	4.0
1 component amounts are expressed as percent of final composition	15 are ex	pressed as	percent of 1	final compo	sition					

N-methyl, N.N-di@-partially hydrogenated talloyloxyethyll, N.Ahydroxyethyl ammonium methyl sulfate (75% in propylane glycol) 3 N-methyl, N.N-diß-partially hydrogenated palmitotyloxyathyll, N-β-hydroxy athyl ammonium methylsulfate (75 % in propylens glycol)

# Example 7

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# Preparation of a Hair Conditioner Composition

To a suitable vessel equipped with stirring, heating, and cooling capabilities was added 426g of delonized water, 2.5g of N.N-dif-partially hydrogenated patritior/toxy ethyl), N.P-hydroxyethyl amine, and 11.0g of a 70% mixture of N.H-dif-partially hydrogenated permittry/toxy ethyl, N.P-hydroxyethyl ammo-

<sup>4</sup> emulsion of an amine functional polymer (Dow Corning) 5 microemulsion of an amine functional polymer (Dow Corning)

nium metriri sullate in propyane glycot. Nits mituture was stirred and neated to 70-75°C. At 70-75°C, 1 5g of cetyl alcohol verer added and the mixture amulatified for 20-30 minutes at 70-75°C. The mixture was then cooled about 50°C at which there 6.5g of a 20 percent aqueous potassium chiorice (KGI) collution, was added. This mixture was then streed well and 10g of Silicone DC 345, 285 go of Silicone X2-839, and 1.00g of DMDM hydranion were added. The composition, 7A, was strived until homogeneous and the pit was checked and adjusted if necessary to between 4.0 and 4.5 with 50% aqueous cliric acid or 50 percent agreems. NDOH.

Conditioner compositions 78, 70, 70, 7E, 7F, and 7G were propared using essentially the came procudure set forth above for propering composition 7A. Table by Valow's since 9the amount of each 1c component in compositions 7A-7E. Each of these compositions resulted in excellent conditioning of hair as demonstrated in both sation and laboratory setting.

Table V

Component <sup>1</sup>	7A	78	7C	70	7E
Delonized water	85.09	85.79	93.50	93.80	85.09
hydroxyethylcellulose		***		0.70	0.70
palm ester amine <sup>2</sup>	0.50	2.00	2.00	2.00	2.00
palm ester quaternary ammonium sait <sup>3</sup>	2.20	~			
cetyl alcohol	3.00	3.00	3.00	2.00	3.00
20% aqueous KCI	1.30	1.30	1.30	1.30	1.30
Silicone DC 345	2.00	2.00			2.00
Silicone DC X2-8939	5.71	5.71	~		5.71
50% aq. NaOH	Q.S.	0.5.	Q.S.	Q.S.	Q.S.
50% aq citric acid	Q.S.	Q.S.	Q.S.	Q.S.	Q.S.
DMDM hydantoin	0.20	0.20	0.20	0.20	9.20

- 1 component amounts are expressed as percent of final composition
- 2 N,N-di(β-partially hydrogenated palmitoyloxy ethyl), N-β-hydroxyethyl amine (neat)
  - 3 N-methyl, N,N-di(g-partially hydrogenated palmito/loxy ethyl), N-g-hydroxy ethyl ammonium methyl sulfate (75% in propylene glycol)

From the foregoing it will be appreciated that, although specific embodiments of the invention have so been described herein for purposes of illustration, various modifications may be made without devisating from the spirit and scope of the invention.

#### Claims

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- 40 1. A hair conditioning composition comprising an effective conditioning amount of an alkanolamine derivative selected from the group consisting of:
  - an alkanolamine ester of the formula:

where

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- A and B are are the same or different and represent lower alkylene having 1-6 carbon atoms; and R<sub>1</sub> and R<sub>2</sub> are the same or different and represent alkyl or alkenyl groups having 8 to 40 carbon atoms;
  - an alkanotamine ester quaternary ammonium salt of the formula:

where

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R is lower alkyl having 1-6 carbon atoms;

A and B are the same or different and represent lower alkylene having 1-6 carbon atoms; and

R<sub>1</sub> and R<sub>2</sub> are the same or different and represent alkyl or elkenyl groups having 8 to 40 carbon atoms; and

X is Ci\*, Br\*, i\*, RSO2\*, or RSO4\*; and

mixtures of the alkanolamine ester and the alkanolamine ester quaternary ammonium sait.

- A conditioning composition according to claim 1, wherein the amount of the alkanolamine derivative in the composition is about 0.1 % to about 20%.
- A conditioning composition according to claim 2, wherein the amount of the alkanolamine derivative in the composition is about 1% to about 10%.
- 25 4. A conditioning composition according to claim 3, further comprising a silicone oil conditioning compound.
  - A conditioning composition according to claim 4, wherein the amount of the silicone conditioning compound is about 0.1% to about 25%.
  - A conditioning composition according to claim 5, wherein the amount of the silicone conditioning compound is about 0.5% to about 5.0%.
  - A composition according to claim 6, wherein the alkanolamine derivative is an alkanolamine derivative queternary ammonium salt and R is methyt, R, and R<sub>2</sub> each is a mixture of C<sub>1</sub>+b<sub>2</sub> and C<sub>1</sub>+b<sub>2</sub> alkyl radicals, A and B are the same and represent ethylene, and X is methyl sulfate.
  - A composition according to claim 6, wherein the alkanolamine derivative is an alkanolamine derivative quaternary ammonium salt end fi is methyl, R; and R; each is a mixture of C;rHs; and C;rHs; alkyl radicals. A and B are the same and represent ethylene, and X is methy sultate.
  - A composition eccording to claim 6, wherein the alkanolamine derivative is an alkanolamine aster and Ri and Ri<sub>2</sub> each is a mixture of C<sub>15</sub>H<sub>27</sub> and C<sub>15</sub>H<sub>29</sub> alkyl radicals, and A and B are the same and represent chilylene.
  - A composition according to claim 6, wherein the alkanolamina derivative is an alkanolamine ester and Ri and R<sub>2</sub> each is a mixture of C<sub>17</sub>H<sub>31</sub> and C<sub>17</sub>H<sub>32</sub> alkyl radicals, and A and B are the same and represent eithylene.
- A composition according to claim 6, wherein the alkanolamine derivative is N-methyl, N,N-di(β-partially hydrogenated talloyloxyethyl), N-β-hydroxyethyl ammonium methyl sulfate.
  - 12. A composition according to claim 6, wherein the alkanotamine derivative is N-Methyl, N,N-di(β-partially hydrogenated palmitoyloxy ethyl), N-β-hydroxy ethyl ammonium methyl sulfate.
  - A composition according to claim 6, wherein the alkanolamine derivative is N-Methyl, N,N-di(β-partially hydrogenated palmitoyloxy ethyl), N-β-hydroxy ethyl ammonium methyl sulfate.

- 14. A composition according to claim 3, wherein the silicone conditioning compound is selected from the group consisting of amodimethicones, dimethicones, cyclomethicones, and mixtures thereof.
- 15. A method for preparing a hair conditioning composition comprising the steps of:
  - (a) mixing water with an emulsifier;(b) heating the mixture;
    - (c) adding an alkanolamine derivative selected from the group consisting of:
      - an alkanolamine ester of the formula:

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A and B are are the same or different and represent lower alkylene having 1-6 carbon atoms; R<sub>1</sub> and R<sub>2</sub> are the same or different and represent alkyl or alkentyl groups having 8 to 40 carbon.

an alkanolamine ester quaternary ammonium salt of the formula:

where

R is lower alkyl having 1-8 carbon atoms;

A and B are the same or different and represent lower alkylene having 1-6 carbon atoms; and R<sub>1</sub> and R<sub>2</sub> are the same or different and represent alkyl or alkenyl groups having 8 to 40 carbon atoms; and

X is Cir., Brr., Ir., RSO<sub>3</sub>r., or RSO<sub>4</sub>r; and

mixtures of the alkanolamine ester and the alkanolamine derivative quaternary ammonium salt;

(d) cooling the mixture.

- 16. A method according to Claim 15, wherein the alkanolamine derivative is added as a mixture of the alkanolamine derivative in an alcohol or glycol.
- 17. A method according to Claim 18, wherein the alkanolamine derivative is added as a mixture of the alkanolamine derivative in isopropanoi, ethenol or propylene glycol.
  - 18. A method of conditioning heir comprising treating the hair with a conditioning composition comprising an alkanolamine derivative selected from the group consisting of:
    - an alkanolamine ester of the formula:

where

A and B are are the same or different and represent lower alkylene having 1-6 carbon atoms;

R<sub>1</sub> and R<sub>2</sub> are the same or different and represent alkyl or alkenyl groups having 8 to 40 carbon atoms:

an alkanolamine ester quaternary ammonium salt of the formula:

$$\begin{array}{c} & \circ \\ & \circ \\ & | | \\ A - OC - R, \\ & + OC - R_2 \end{array}$$

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where R is lower alkyl having 1-6 carbon atoms;

A and B are the same or different and represent lower alkylene having 1-6 carbon atoms; and

R<sub>1</sub> and R<sub>2</sub> are the same or different and represent alkyl or alkenyl groups having 8 to 40 carbon atoms; and

X is Cir., Br., Ir., RSOs., or RSOs.; and

mixtures of the alkanotamine ester and the alkanotamine derivative quaternary ammonium salt.

- 35 19. A method according to Claim 18, wherein the composition further comprises a silicone conditioning compound.
  - 20. A conditioning composition according to Claim 1, whareing the alkanolamine derivative is a mixture of alkanolamine seter and alkanolamine ester quaternary ammonium sait, having a ratio of alkanolamine ester optimization of alkanolamine and alkanolamine armonium sait of about 1:50.



#### EUROPEAN SEARCH REPORT

EP 93 42 0326

Category	Citation of document witi of relevant	a indication, where appropriate, passages	Reservant to claims	CLASSIFICATION OF THE APPLICATION (6st. CL5)
Χ.	DE-A-4 138 630 (HE	ENKEL KGAA ET AL)	1-3, 11-13, 15-18	A61K7/06 A61K7/50 A61K7/48
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	* page 11, line 20 * claims 1-4,6-7,1	- line 47 *		
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